

1 Ecological niche overlap between co-occurring native and exotic ungulates:
2 insights for a conservation conflict

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14 **Abstract**

15 Exploitative competition implies an indirect interaction in which a resource exploited by
16 one species is not available for another; e.g., when species share diet or habitat. It plays
17 a key role in community structure and dynamics. Here we evaluated the niche overlap
18 between the exotic aoudad (*Ammotragus lervia*) and the native Iberian ibex (*Capra*
19 *pyrenaica*) where the species coexist in the Iberian Peninsula, along two main
20 dimensions, the trophic niche and the environmental niche. Then we assessed the spatial
21 segregation of the species. We expected that if a niche overlap was high, competition
22 could drive spatial segregation to allow co-existence. We analyzed their trophic niche
23 overlap by using the content of stable isotopes $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ in the hair of both species.
24 To establish environmental niche competition, we compared the similarity in their
25 habitat, estimated by environmental niche models based on the fine-scale presence

records of each species obtained from field surveys. To test if spatial segregation occurred, we analyzed both species' co-occurrence. Our results indicated that both species shared a similar trophic niche measured by stable isotopes, both species showed a similar distribution of suitable areas, and that both species' environmental niches were more similar than expected. Finally, a negative spatial association was found between the aoudad and Iberian ibex. These results reveal that both species are ecologically similar and suggest that fine-scale spatial segregation might have favoured their co-existence in semiarid Mediterranean mountains. Our results show that integrating information on trophic and environmental niche overlap with fine scale spatial distribution might improve the study of competitive interactions among wild ungulates.

Key words: *Ammotragus*; Assisted colonization; *Capra*; Competition; Environmental model; Stable isotopes

1. Introduction

Interspecific interactions such as competition play a key role in the ecosystem structure and dynamics (Barbosa and Castellanos, 2005; Tilman, 1987). For the competition process to happen, the following conditions must be met: 1) different species must share resources; 2) resources must be limited; 3) the joint exploitation of these resources must negatively affect both species' performance (Milne, 1961; Prins, 2000). Competition can be based on interference or exploitation among sympatric species (Begon et al., 2006). Interference competition happens when both species directly interact, preventing another species from exploiting resources (Begon et al., 2006; Linnell and Strand, 2000). Exploitative competition implies indirect interactions in which a resource consumed by one species is not available for another (Lang and Benbow, 2013; de Boer and Prins, 1990).

50 Studies that focus on competition between species that exploit the same resources (i.e.
51 intraguild competition) usually encompass ecological processes with species that have
52 co-evolved (Ballejo et al., 2018; Grassel et al., 2015; Wright 2002). Co-evolved
53 competing species can co-exist because they differ in the realized niche; i.e. resources
54 and conditions that a species exploits as a result of interactions with other species
55 (Giller 1984). According to Hutchinson (1957), the niche concept is defined as an *n*-
56 *hyperdimensional volume*, where *n* is the number of dimensions that compose the niche.
57 Therefore, niche differentiation can occur along different dimensions, such as food or
58 habitat. Among the mechanisms that facilitate co-existence and alleviate competition,
59 predation (Chesson 2000) or spatial segregation on fine scales to exploit shared
60 resources are highlighted processes (e.g. Barrio and Hik, 2013).

61 Novel intraguild competition processes may appear when exotic species are introduced
62 beyond their natural range and interact with the native species they share resources with
63 (Mooney and Cleland 2001). These new interactions can reduce the abundance and
64 richness of native species (Blackburn et al., 2004; Gaertner et al., 2009). Introduced
65 invasive species are considered the second cause of biodiversity loss (Wilcove et al.,
66 1998) and they are associated with alterations of ecosystem processes (Raizada et al.,
67 2008) and the community structure (Hejda et al., 2009). It has also been detected that
68 exotic species may affect the genetic diversity and the evolutionary pathway of native
69 species (Mooney and Cleland 2001). This phenomenon has been evaluated in different
70 areas of the world; e.g., introduced carnivores into Australia (Doherty et al., 2017) or
71 Europe (Bonesi et al., 2004; Harrington et al., 2009), or between exotic and native deer
72 in North America (Faas and Weckerly 2010).

73 In the case of wild ungulates, introductions usually respond to sport hunting interests
74 (Spear and Chown 2009). Different studies on the competition between exotic and

75 native ungulates have concluded that non native species impair native species (Stewart
76 et al., 2002; Dolman and Wäber 2008). Other studies have also shown positive effects,
77 such as facilitation processes between non native and native ungulates. For instance,
78 Gordon (1988) found that during spring the native red deer (*Cervus elaphus*) from the
79 Scottish island of Rum (Pérez-Espona et al. 2013) preferentially grazed in areas that had
80 been grazed by cattle during the previous winter, because cattle improves forage
81 availability for red deer. Another example of facilitation was described by Odadi et al.
82 (2011). These authors detected that during the wet season in a savannah ecosystem, the
83 zebras (*Equus* spp.) decreased cover of dead grass and the cattle got access to higher-
84 quality food. So, the interactions between native and non native ungulates could vary
85 among different systems.

86 Besides translocations, wild ungulates are currently in a process of recolonization in
87 developed countries (Apollonio et al., 2010). This rewilding process leads to an
88 increased spatial overlap among ungulates. Considering that current ecosystems are
89 subject to intense human influence which includes the elimination of predators, the
90 presence of livestock and the introduction of exotic species (Latham 1999) novel
91 competition interactions may arise.

92 Both natural rewilding and introductions are commonplace in Europe, including
93 Mediterranean ecosystems (Apollonio et al., 2010). The Iberian ibex, an endemic
94 ungulate to the Iberian Peninsula, had disappeared in most of its range. Nevertheless,
95 since the mid-twentieth century, its populations have increased and large areas from
96 which it had disappeared have been recolonised (Acevedo and Cassinello 2009).

97 The aoudad is an ungulate native from North Africa, whose populations are decreasing
98 in its native range due to habitat loss or degradation and human persecution (Durant et

99 al., 2014). Currently the species is catalogued as vulnerable by IUCN in its native range
100 (Cassinello et al., 2008). However, it has been introduced into several countries of
101 Europe and America in the twentieth century. In southeastern Spain the aoudad was
102 introduced in the 1970s motivated by hunting interests (Cassinello 1998, Valverde
103 2005) to protect the semiarid landscapes of Murcia through the presence of a hunting
104 large herbivore adapted to the dry conditions (Valverde 2005) and probably also for
105 conservation purpose (Valverde 2004). The individuals for the introduction came from
106 different zoos and from unknown African origin. The fact of the introduction of an
107 exotic species outside of its native range for conservation purpose is currently known as
108 “assisted colonization” (Seddon 2010). Thus, the introduction of the aoudad in the
109 Iberian peninsula for conservation purpose might be considered a pioneering example of
110 an assisted colonization before this concept had been fully established. Moreover,
111 animals from the Western Sahara (Spanish protectorate at that time) of several threaten
112 ungulates (dama gazelle *Nanger dama*, dorcas gazelle *Gazella dorcas*, Cuvier’s gazelle
113 *Gazella cuvieri*, and aoudad) were brought to a captive breeding center created in Spain
114 in 1971 for conservation purposes (currently called Experimental Station of Arid Zones,
115 EEZA by its initials in Spanish; <http://www.eeza.csic.es>).

116 Since the 1990s, both the aoudad and Iberian ibex have co-existed in some mountains of
117 the Region of Murcia (SE Spain), and the expansion of both species has been apparently
118 influenced by habitat connectivity and interspecific competition (Anadón et al., 2018).
119 Previous studies indicate that the introduced aoudad could compete with native
120 ungulates, particularly with the Iberian ibex (Acevedo et al., 2007; but see Cassinello
121 2018). Besides the aoudad has been described as a potential hazard to threatened
122 vegetation due to high population densities, although this deleterious effect could be
123 caused by other overabundant large herbivores, even native species (Velamazán et al.,

2017). Furthermore, crop damage has been also pointed out as another emergent issue leading to the application of management tools such as diversionary feeding to mitigate these impacts (Pascual-Rico et al., 2018). Due to these problems associated with the exotic species (see Mori et al. 2017), the aoudad was first included in the Spanish catalogue of invasive species in 2013 (Real Decreto 630/2013, Spanish Government) although the population in Murcia Region introduced legally before 2007 was excluded. However, in a sentence by the Spanish Supreme Court (sentence 637/2016) following a demand by conservationist, the aoudad was included as an invasive exotic in all its range within the Spanish territory, which spurred a heated debate among the main stakeholders (i.e. wildlife managers, hunters, ecologist, farmers and conservationists). The potential competition between the aoudad and the Iberian Ibex was one of the main arguments to consider the exotic aoudad as an invasive species. Nevertheless, after national Government approved a law (Law 7/2018) to return to the situation in which aoudad populations introduced legally before 2007 are not considered as invasive species. Given the changing legislative situation of a current conservation conflict among the stakeholders abovementioned, it is necessary to assess with scientific criteria the potential competition between the aoudad and the Iberian ibex.

Our main objective with the present study was to evaluate the potential competition between the Iberian ibex (native species) and the aoudad (exotic species) in the mountain ranges where they live in sympatry. Specifically, we evaluated the ecological niche overlap along two main dimensions, the environmental niche and the trophic niche, because they are the commonest partitioned dimensions (Schoener 1983; Toft 1985). Then we assessed the spatial co-occurrence of both species on the fine scale. We expected that if a niche overlap was high, competition could drive spatial segregation to allow co-existence.

149 **2. Material and methods**

150 **2.1 Study area**

151 The study was conducted in different areas in the Region of Murcia (SE Iberian
152 Peninsula; see Figure 1 and Table A.1) where both the Iberian ibex and aoudad
153 (Artiodactyla order) co-exist. The study area includes mountain ranges with a mean of
154 $116.7 \pm 71.6 \text{ km}^2$ and $1239.6 \pm 321.1 \text{ m.a.s.l.}$ (see Table A.1 for more detailed data on
155 each mountain range included in the study). The climate is Mediterranean, the mean
156 temperature range from 10° to 16°C , and the mean rainfall of 375 mm per year.

157 Region of Murcia forms part of the Iberian ibex's historical range, which drastically
158 declined in the first half of the twentieth century, followed by recovery over the last five
159 decades (Anadón et al., 2018). The aoudad was introduced into the region in the 1970s
160 for hunting purposes, specifically in the Sierra Espuña Regional Park ($37^\circ 47' - 37^\circ 56' \text{N}$,
161 $1^\circ 27' - 1^\circ 40' \text{W}$). Since then, this exotic species has spread to other mountains in the
162 region (see Supplementary Material A.1 for more biological information about study
163 species).

164 **2.2 Trophic niche overlap**

165 We, in collaboration with local hunters, collected hair samples of both ungulates
166 (Iberian ibex $n=25$, of which 10 males, 13 females and 2 indeterminate samples; aoudad
167 $n=26$, of which 13 males and 13 females) during different hunting seasons in 2013 and
168 2014 in the Region of Murcia (Figure 1). In all cases, the frontal dorsal hair nearest the
169 skin was collected and cut by stainless steel surgical scissors. All hair samples were
170 immediately stored in a paper envelope tube until their isotopic analyses were
171 conducted. All hair samples were cleaned and powdered, and 0.3-0.4 mg of each hair

sample was packed into tin capsules. Isotopic analyses were performed at the Laboratory of Stable Isotopes of the Estación Biológica de Doñana (www.ebd.csic.es/lie/index.html). Samples were combusted at 1020°C using a continuous flow isotope-ratio mass spectrometry system (Thermo Electron) by means of a Flash HT Plus elemental analyzer interfaced with a Delta V Advantage mass spectrometer which applies international standards, run each 9 samples; LIE-CV and LIE-PA, previously normalized with the international standards IAEA-CH-3, IAEA-CH-6, IAEA-N-1 and IAEA-N-2. Stable isotope ratios were expressed in the standard δ -notation (‰) relative to Vienna Pee Dee Belemnite ($\delta^{13}\text{C}$) and atmospheric N_2 ($\delta^{15}\text{N}$). Based on laboratory standards, the measurement error (standard deviation) was ± 0.1 and ± 0.2 for $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$, respectively.

The trophic niche overlap was measured by comparing the degree of isotopic overlap between species (Jackson et al., 2011). As alternative to the isotopic metrics provided in SIBER procedures (Jackson et al., 2011), we examined the isotopic niche overlap between species adapting the framework proposed by Broennimann et al., (2012), which applies kernel smoothers to species occurrence in a two-dimensional gridded space. In our case, this space was defined for the respective overlap analysis by the $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values of all the individuals (see Navarro et al., 2015). Specifically, the trophic overlap based on the isotopic values of both species was calculated using the D-metric, which ranges from 0 (no overlap) to 1 (complete overlap). We applied a permutation-based approach (100 permutations) to evaluate whether the overlap values were higher than expected at random according to the available isotopic space (similarity test, Warren et al., 2008). These analyses were conducted using the ‘ecospat’ library in R software. Interspecific differences in the $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values were tested by using Student’s t-tests. The significance level for all the tests was set at $p < 0.05$.

198 2.3 Environmental niche overlap

199 We developed an environmental niche model (ENM) for both species in the mountains
200 that they share. Then we compared the similarity of the resulting models. We used the
201 Maximum Entropy Software, Maxent 3.3.3 k (Phillips and Dudik 2008), to develop an
202 ENM of the two species independently using presence-only data. We compiled presence
203 records of each species using the observations obtained from the autumn censuses of
204 target species. These censuses were made in 2012, 2014, 2015 and 2016 in the
205 mountains where both species were present (Iberian ibex n=174; aoudad n=429).
206 Species position data were transferred to a geographic information system. We assigned
207 the presence records to a grid cell matrix of 200 x 200 m of the study area. We selected
208 this grid size to obtain the highest resolution of the study area and to avoid the sampling
209 error when assigning the field observations in the map. We used the area under the
210 curve (AUC) that derived from the test data as a yardstick provided by Maxent to
211 evaluate the model's efficacy. We considered test AUC values > 0.75 with sufficient
212 discriminatory capacity (Elith 2000).

213 To predict the habitat suitability of both the Iberian ibex and aoudad, we used land
214 cover and topography variables as environmental predictors (Table 1). Land cover
215 variables represented the percentage cover of the habitat in the study area in a 200x200
216 m grid cell. Topography variables were obtained from a 5-meter resolution digital
217 elevation model, from which the elevation and slope variables were developed. The
218 nature of our variables as well as the size of extent and grain of our models thus define
219 niche overlap analysis related to landscape habitat use patterns, rather than distribution
220 limiting factors, that would act at larger spatial scales.

We employed ENMToolsv.1.4.4 (Warren et al., 2010) to measure the similarity of the ENMs generated with MaxEnt. With this software, we calculated Schoener's (1968) D index to quantify niche similarity, which was estimated by comparing habitat suitability for each grid cell of the study area using ENMs.

To test whether ENMs were more similar than expected by chance, we ran a background test. To do so, the test generates a null distribution for the ENM difference expected between one species and the occurrence points placed at random within the range of the other species. The niche similarity hypothesis among species is rejected if Schoener's D is significantly higher or lower than those expected from the null distribution (Warren et al., 2010). In our case, significantly higher or lower values implies that D is over the 0.975 percentile or below the 0.025 percentile (the equivalent to $p = 0.025$) of the null distribution values (two-tailed comparison), respectively.

2.4 Spatial segregation

We tested if our study species more or less co-occurred spatially than expected by random on the 200 x 200m grid scale. To calculate the number of cells occupied by at least one species, we used QGIS (2017). The obtained dataset consisted in a presence-absence matrix with rows taken as species ($n=2$) and columns as occupied cells ($n=465$). We used the "co-occur" package in R (Griffith et al., 2016), which applies the probabilistic model of species' co-occurrence (Veech 2013) to assess if species co-occur positively or negatively.

3. Results

3.1 Trophic niche overlap

The $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values did not differ statistically between species ($\delta^{15}\text{N}$, Iberian ibex = 3.68 ± 1.05 ‰, aoudad = 3.38 ± 0.95 ‰, $t = -1.08$, $p = 0.28$; $\delta^{13}\text{C}$, Iberian ibex =

-23.85 \pm 0.66 ‰, aoudad = -23.94 \pm 0.87‰, $t = -0.42$, $p = 0.67$). Similarly to the isotopic values, the isotopic niche overlap was high between both species (D-index = 0.59, $p = 0.23$), the trophic niche of the aoudad overlapped 84.4% the trophic niche of the Iberian ibex, and conversely, the trophic niche of the Iberian ibex overlapped 77.6% the trophic niche of the aoudad (Figure 2).

3.2 Environmental niche overlap

The AUC of the MaxEnt model for the Iberian ibex was 0.88, and the AUC for the aoudad was of 0.82. Both species showed a very similar distribution of suitable areas (Figure 3). For the Iberian ibex, the model showed that this species correlated mainly with elevation and slope, and negatively with croplands. The other variables contributed less than 5% to the model. With the aoudad, croplands contributed the most and negatively to the model, as did slope, elevation and grassland (Figure 4).

The environmental niche overlap between the studied species was quantified using Schoener's D index, which was 0.71. The background test indicated that our study species were more similar than expected by chance (Schoener's D > 97.5% of the null distribution values; Figure 5).

3.3 Spatial segregation

In the spatial segregation analysis on the fine scale, of the 465 cells of 200 x 200m occupied by at least one of the two species, we found that only six were both co-occurred species. The spatial co-occurrence analysis showed a negative significant association between the Iberian ibex and the aoudad ($p < 0.001$).

4. Discussion

Our study case provides us with the opportunity to evaluate, in several ways, the differences in the ecological niche between exotic and native species of the same ecological guild. According to the stable isotope analysis results, the Iberian ibex and the aoudad showed high trophic niche similarity. Regarding habitat selection based on environmental niche models, the study species showed a large overlap in habitat use, but with slight differences on the fine scale. In the co-occurrence test, as the studied species showed a negative association, they avoided co-occurring on the fine scale. These results indicate that both species potentially compete in the trophic and environmental niche dimensions, and that spatial segregation might be a key mechanism to allow for long-term co-existence.

4.1 Trophic niche overlap

Although stable isotope analyses do not provide definitive assessment of diet in the manner of stomach content analysis, feeding observations or faecal analysis (Layman et al 2012), it is a useful tool employed for reconstructing diets (Kelly 2000, Layman et al., 2012). The isotopic analysis of hair implies that the levels of nitrogen and carbon correspond to the plants eaten several months before. So, the results reflect the diet of the animals during the spring and early summer, when the plant productivity (NDVI) is maximum in the region (see Barbosa et al. 2019) and therefore the food available is greater than during the rest of the year. $\delta^{15}\text{N}$ is a proxy of a species' trophic level (Peterson and Fry 1987) and, although herbivores can consume vegetation with different $\delta^{15}\text{N}$ levels (Pacyna et al., 2018), no differences were found in our study species. Nevertheless, similar $\delta^{13}\text{C}$ values indicate that species feed on the same kind of vegetation, in our case C3 plants; i.e. they were mainly browsers in our study area. However, it must be considered that the feeding patterns of ungulates do not remain constant over time and some species show high feeding plasticity (Acevedo and

293 Cassinello 2009; Lehmann et al., 2011). Studies about the Iberian ibex have
294 demonstrated that browsing focuses on shrub or tree species. (Martínez 1989; Martínez
295 2002). The aoudad in the southeast of the Iberian Peninsula also showed preference for
296 shrub species (Fernández-Olalla et al., 2016). Moreover, our study species also feed on
297 grass and forb species (Martínez 1989; Martínez 2002; San Miguel et al., 2010).
298 Therefore, the proportion of browse, grass and forb eaten by both the Iberian ibex and
299 the aoudad might depend on vegetation availability, season and the weather conditions
300 (Cassinello 1998; Wilson and Mittermeier 2011). For example in their natural
301 distribution area, both species can habit from sea level up to about 3000 – 4000 m
302 (Cassinello 1998; Granados et al., 2007), which indicates that they can occupy
303 ecosystems with different plant communities. Moreover, the dietary plasticity of both
304 ungulates allows them to feed on less palatable plants, such as *Rhamnus lycioides*
305 bushes and *Pinus halepensis* trees (San Miguel et al., 2010). In our case, and according
306 to the values established by Moreno-Gutiérrez et al., (2012) for $\delta^{13}\text{C}$ of leaf cellulose for
307 several plant species, *S. tenacissima* forbs and *R. lycioides* bushes may form an
308 important part of the diet of both the Iberian ibex and the aoudad in our study area.

309 The aoudad showed a higher standard deviation of the $\delta^{13}\text{C}$ levels than the Iberian ibex.
310 Hence its trophic amplitude was wider. The trophic niche overlap of the aoudad on the
311 Iberian ibex was higher than that of the Iberian ibex on the aoudad. These results may
312 indicate that the aoudad feeds on a higher diversity of plants than the Iberian ibex,
313 which could be interpreted as an advantage for the exotic species *versus* the native
314 species. Nevertheless, native herbivores may have narrower trophic niches than
315 sympatric exotic species because the diet of the former includes fewer items, but
316 exploits better the resource by feeding on the most nutritious plants available (Jarman
317 and Sinclair 1979; Reus et al., 2017).

318 **4.2 Environmental niche overlap**

319 The ecological niche models reveal that both the Iberian ibex and the aoudad similarly
320 respond to habitat features, although the percentage contribution of each variable varied.
321 The distribution of both species is positively linked to elevation and slope, and altered
322 areas (i.e. croplands) appeared to be avoided. It is interesting that despite including only
323 the mountain ranges where both species cohabit in this study, the ENMs slightly
324 differed from one another. This means that the species in these restricted areas did not
325 distribute in the same way. These results agree with previous studies conducted for both
326 species (Acevedo et al., 2007; Anadón et al., 2018).

327 One of the ways to detect ecological niche differences is by comparing the ENMs
328 developed by the MaxEnt software, which has been demonstrated as the most capable
329 method for modelling distributions of mammals and other species (Hernandez et al.,
330 2008; Phillips et al., 2006). According to de Boer and Prins (1990), a large overlap in
331 habitat use could be a sign of non problematic co-existence between two species, while
332 a small overlap might indicate segregation processes due to competition. The niche
333 overlap found for our study species was large (Schoener's D index=0.71), and the
334 background test showed that the environmental niches for both species were more
335 similar than expected. This may be related to the fact that both species (included in the
336 Caprini tribe) present morphological, biological and behavioural similarities.

337 **4.3 Spatial segregation**

338 If the niche overlap was large, we expected competition possibly drive spatial
339 segregation. Our results showed that spatial segregation could already be acting as a
340 mechanism to allow for co-existence, despite the interaction noted between our study
341 species being a result of recent human intervention (with no common evolutionary
342 history). Therefore, if we consider that our results indicate that both species are

ecologically similar in terms of the evaluated niche dimensions and the detected spatial segregation, then both species could compete, especially when resources are limited; e.g., when environmental perturbations occur, such as drought periods. Competition between both the Iberian ibex and the aoudad could lead to one species' displacement, and even to one of them eliminating the other through the principle of competitive exclusion; i.e. one competing species eliminating or excluding another species (Hardin 1960). To determine the output (i.e. competitive exclusion or co-existence), it will be necessary to estimate the competition coefficient of each species and the carrying capacity of the environment in future research.

In summary, in the given conditions, the strong similarity in the trophic and environmental niche of both the Iberian ibex and the aoudad indicates potential competition between them. However, the spatial segregation on the fine scale seems to act as a mechanism to facilitate the co-existence between the native and exotic ungulate species.

Currently the aoudad has been eliminated of the Spanish Catalogue of Invasive Exotic Species for some areas where it was introduced. However, Carboneras et al., (2017) created a prioritised list of invasive alien species where the species are ranked according to their potential threat to biodiversity in Europe. According to this list, the aoudad is considered in an invasion phase and it may be a major impact to biodiversity and ecosystems. This study sheds light on the research gap that exists on the interaction between the Iberian ibex and the exotic aoudad, which can be applied in the development of wildlife policies aimed at better management and conservation of the species.

367 The zoologist J. A. Valverde anticipated the future consequences of human activities on
368 the Sahara's megafauna (see Durant et al., 2014; Brito et al., 2018) when he proposed
369 the aoudad introduction in the Iberian Peninsula (Valverde 2004). However, this
370 pioneering assisted colonization example, accomplished more than 40 years ago, failed
371 to forecast the ongoing conflicts among different stakeholders and native species from a
372 conservation perspective. Nowadays, assisted colonization is a controversial tool due to
373 the possible consequences that may result (Hoegh-Guldberg et al., 2008; Ricciardi and
374 Simberloff 2009), both short and long-term. Although it is rarely used (e.g. Kuussaari et
375 al., 2011), there are several proposals for its application (e.g.
376 <https://theaustralianrhinoproject.org>) that should be done with caution, if finally this
377 tool is applied (Loss et al., 2011).

378 The situation of the aoudad in Spain is paradoxical, since it is catalogued as vulnerable
379 in its native range. However, there are other similar cases around the world. The
380 Philippine deer (*Rusa marianna*) and the banteng (*Bos javanicus*) are vulnerable and
381 endangered, respectively, in their native range. However, these two species were
382 introduced in other non-native areas where they established in the wild (Bradshaw et al.,
383 2006; MacKinnon et al. 2015). Gibson and Yong (2017) proposed translocation of these
384 introduced populations to their native ranges if threats to the species disappear. Thus, it
385 is possible to mitigate conflicts and impacts in the non-native areas and contribute to the
386 conservation of species in their native range.

387 Ungulates (Artiodactyla order) are the mammals with the highest proportion of
388 successful introductions around the world (Clout and Russell 2007). In their natural
389 ranges, their abundance and distribution are increasing, which also occurs in new
390 colonised areas for introduced species (Apollonio et al., 2010). The consequences of
391 herbivore ungulate introduction, whatever the reason, can alter biodiversity interactions

(Vázquez and Simberloff 2003), and tend to strongly impact the new ecosystems that they occupy (Duffy 2003). One of the new processes that may appear is competition with native species, especially within the same guild (see Dolman and Wäber 2007). However, competition interactions are not easy to demonstrate in the field because manipulations to evaluate changes in carrying capacity and population dynamics in relation to the relative abundance of interacting species are difficult to perform in the wild (Hakkarainen and Korpimäki 1996). Nevertheless, our results show that integrating information on trophic and environmental niche overlap with fine scale spatial distribution might improve the study of competitive interactions among wild ungulates and support science-based management decisions.

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611 **Figures and Tables**

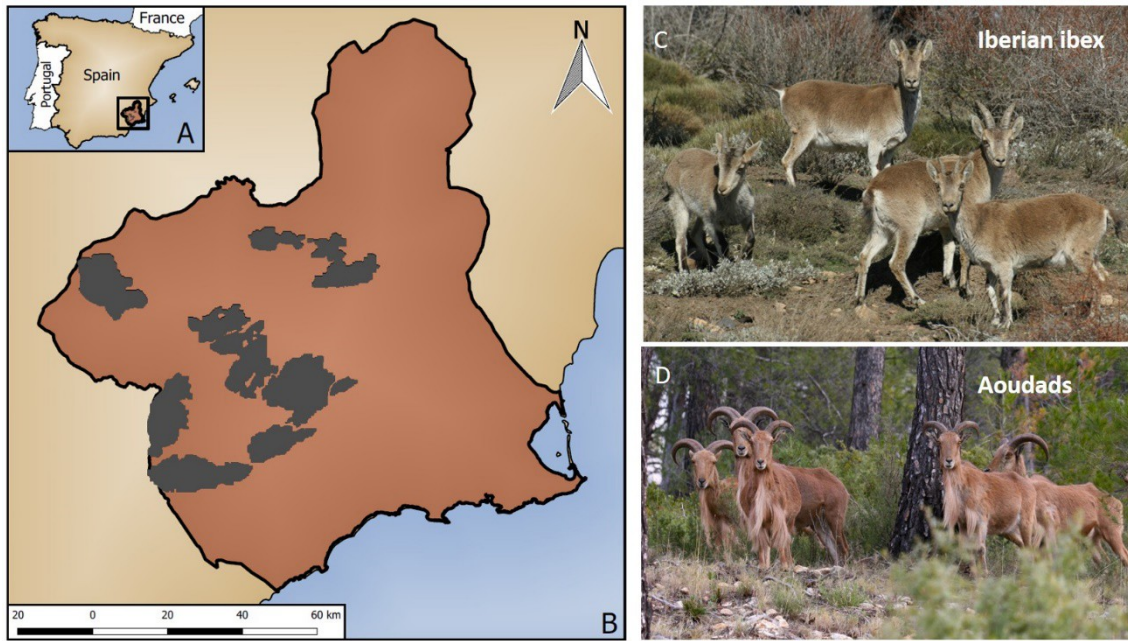
612 Figure 1. (A) Position of the study area in the Iberian Peninsula. (B) Map of the study
613 area (mountains in the Region of Murcia, SE Spain). Black areas indicate the mountains
614 where the Iberian ibex and the aoudad co-exist. (C) and (D) are pictures of the study
615 species in the study area.

616 Figure 2. Isotopic niche of both the Iberian ibex and the aoudad, and the overlap niche
617 between them.

618 Figure 3. Habitat suitability models for both the aoudad and Iberian ibex as assessed
619 from MaxEnt. The province and the Sierra Espuña Regional Park limits (black lines) are
620 shown for spatial reference.

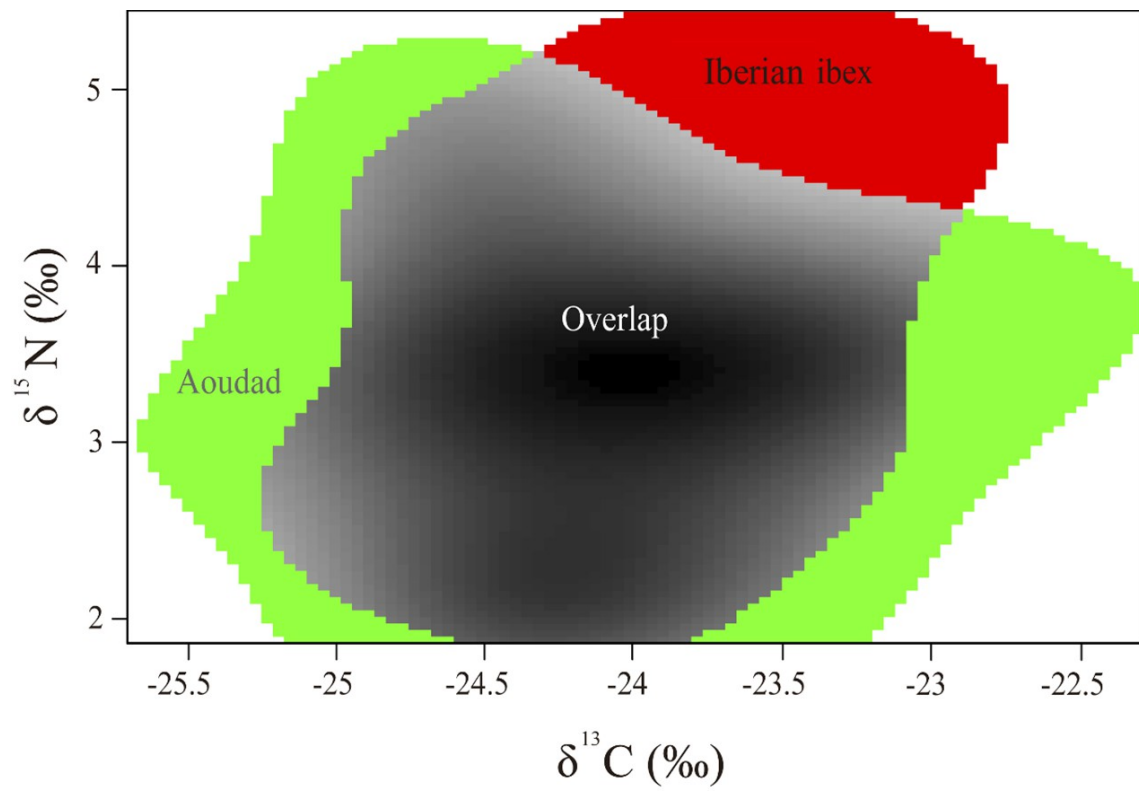
621 Figure 4. Contribution of the environmental variables to construct the MaxEnt
622 environmental niche models for both the Iberian ibex (blue bars) and aoudad (orange
623 bars).

624 Figure 5. Background test histograms for the Iberian ibex (blue bars) and the aoudad
625 (orange bars). Schoener's D index (red arrow) was higher for the null distributions
626 generated of both the aoudad and Iberian ibex. This indicates that the two species are
627 more similar than expected based on available habitat.



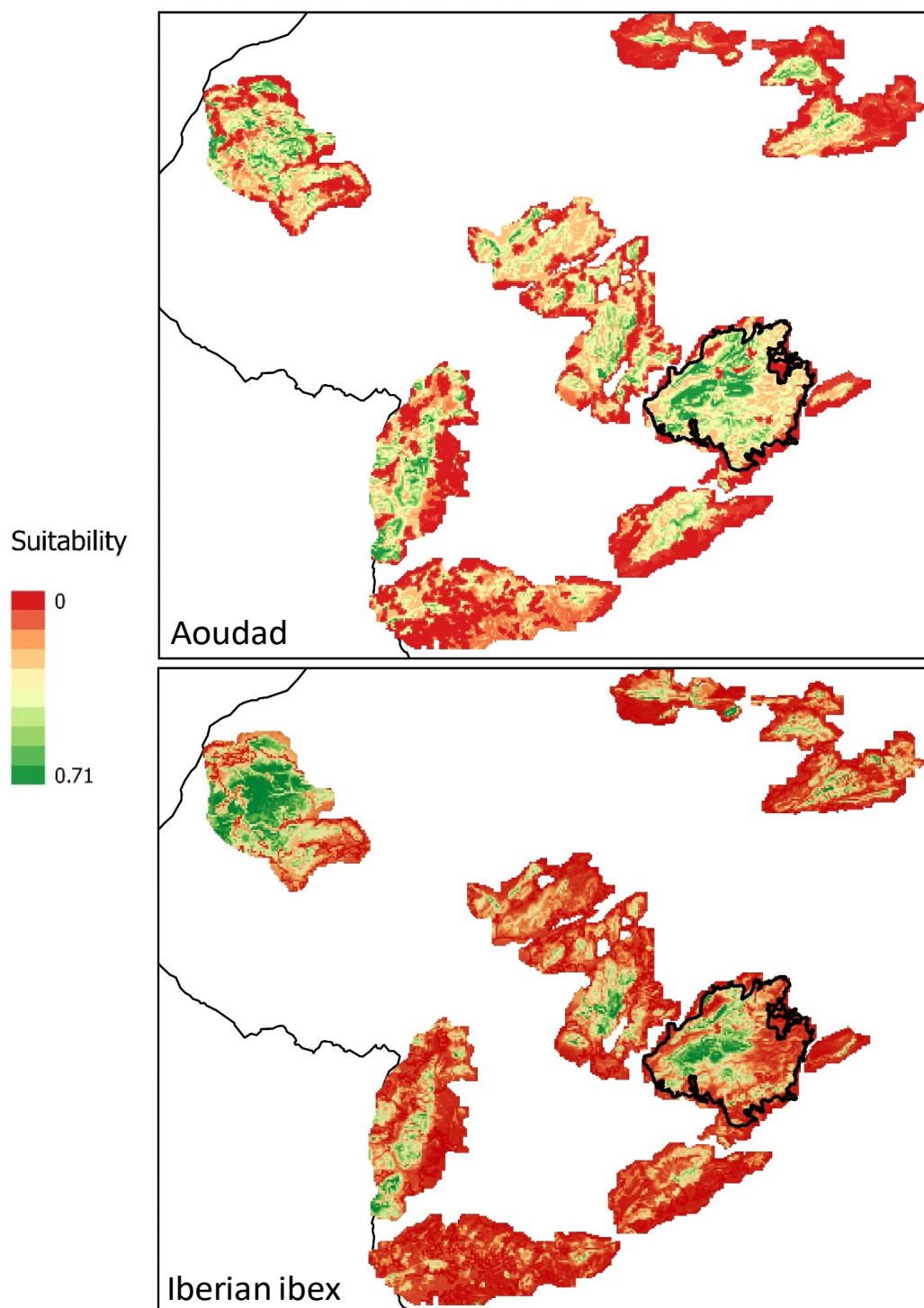
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629 Figure 1



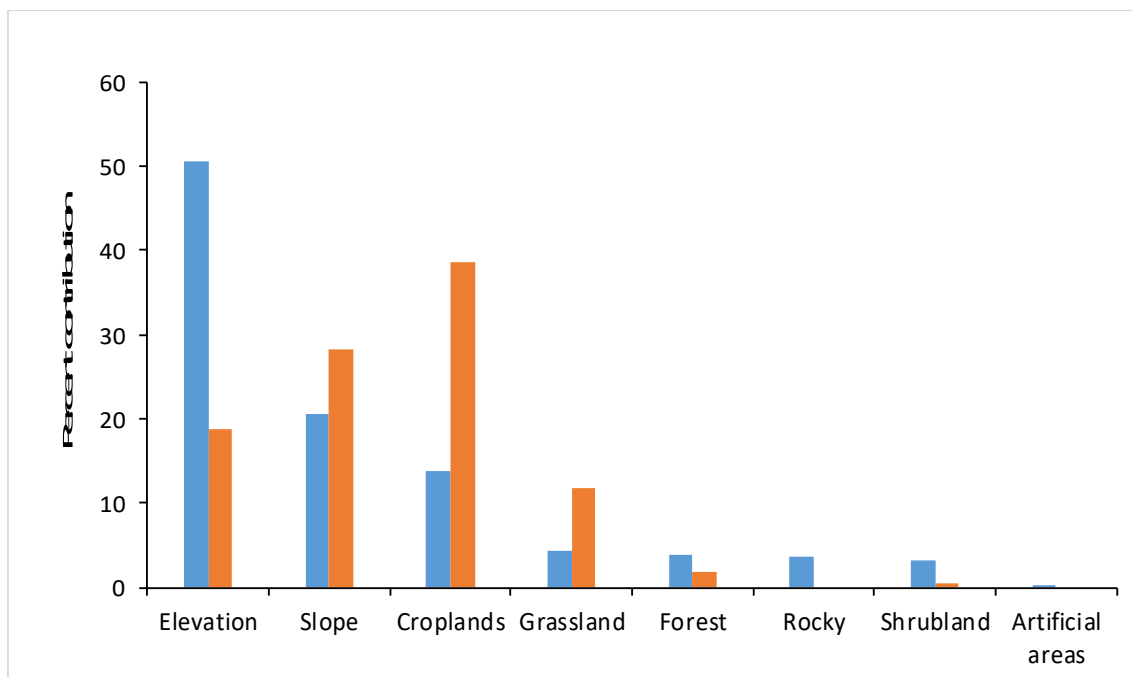
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631 Figure 2



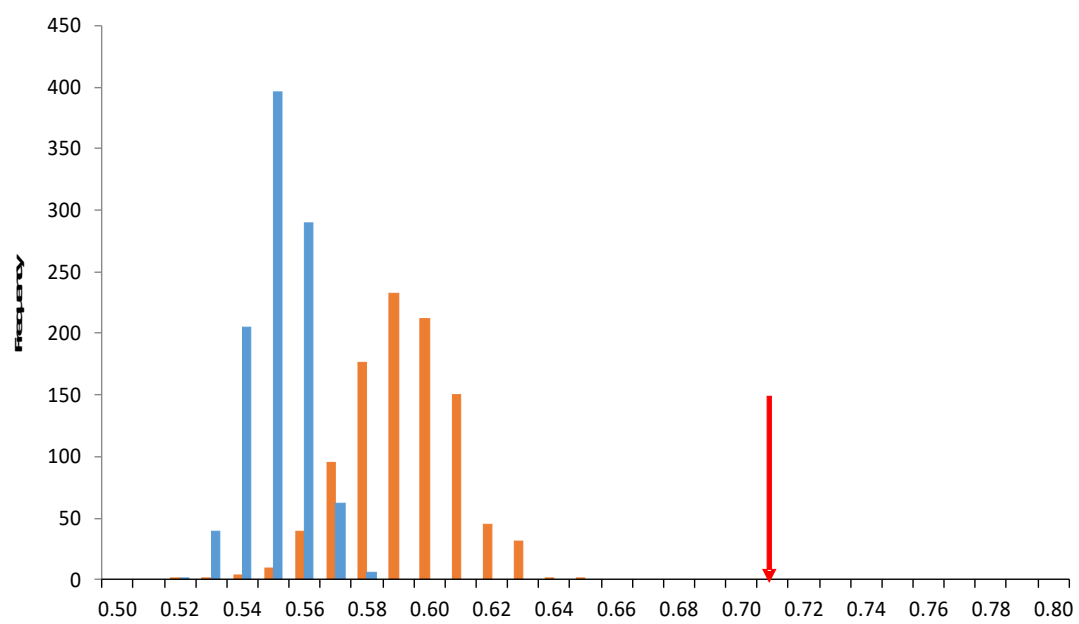
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633 Figure 3



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635 Figure 4



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637 Figure 5

638 **Table 1.** Predictor variables used to assess habitat. All the variables were continuous.

Variable		Source
Land cover	Forest	CLC 2012 CC-BY 4.0 ign.es
	Shrubland	
	Grassland	
	Rocky	
	Croplands	
	Artificial areas	
Topography	Elevation	MDT5 2009 CC-BY 4.0 ign.es
	Slope	Derived from MDT5 2009 CC-BY 4.0 ign.es

639